

1 **Understanding the anthropogenic influence on formation of biogenic**
2 **secondary organic aerosols in Denmark via analysis of organosulfates**
3 **and related oxidation products**

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15

16 **Abstract**

17 Anthropogenic emissions of sulfur dioxide (SO₂) and nitrogen oxides (NO_x) may affect
18 concentration levels and composition of biogenic secondary organic aerosols (BSOA) through
19 photochemical reactions with biogenic organic precursors to form organosulfates and nitrooxy
20 organosulfates. We investigated this influence in a field study from May 19 - June 22, 2011 at two
21 sampling sites in Denmark. Within the study, we identified a substantial number of organic acids,
22 organosulfates and nitrooxy organosulfates in the ambient urban curbside and semi-rural
23 background air. A high degree of correlation in concentrations was found among a group of specific
24 organic acids, organosulfates and nitrooxy organosulfates, which may originate from various
25 precursors, suggesting a common mechanism or factor affecting their concentration levels at the
26 sites. It was proposed that the formation of those species most likely occurred on a larger spatial
27 scale with the compounds being long-range transported to the sites on the days with highest

1 concentrations. The origin of the long-range transported aerosols was investigated using the Hybrid
2 Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model in addition to modeled
3 emissions of related precursors including isoprene and monoterpenes using the global Model of
4 Emissions of Gases and Aerosols from Nature (MEGAN) and SO₂ emissions using the European
5 Monitoring and Evaluation Program (EMEP) database. The local impacts were also studied by
6 examining the correlation between selected species which showed significantly enhanced
7 concentrations at the urban curbside site and the local concentrations of various gases including
8 SO₂, ozone (O₃), NO_x, aerosol acidity and other meteorological conditions. This investigation
9 showed that an inter-play of the local parameters such as the aerosol acidity, NO_x, SO₂, relative
10 humidity (RH), temperature and global radiation seemed to affect the concentration level of those
11 species, suggesting the influence of aqueous aerosol chemistry. The local impacts however seemed
12 minor compared to the regional impacts. The total concentrations of organosulfates and nitrooxy
13 organosulfates contributed to approximately 0.5 - 0.8 % of PM₁ mass on average at the two
14 sampling sites.

15

16 **1 Introduction**

17 Volatile organic compounds (VOC) are emitted from both biogenic and anthropogenic sources and
18 are oxidized to form lower volatility products partitioning between the gas and particle phase,
19 leading to the formation of secondary organic aerosols (SOA) (Donahue et al., 2006; Kroll and
20 Seinfeld, 2008; Jimenez et al., 2009; Hallquist et al., 2009). In general, the production of biogenic
21 SOA (BSOA) is estimated to be significantly larger than anthropogenic SOA (ASOA) (Goldstein
22 and Galbally, 2007; Heald et al., 2010; Spracklen et al., 2011). For example, Hallquist et al. (2009)
23 estimated a many-fold higher annual global production of BSOA (88 TgC) compared to ASOA (10
24 TgC), which were however associated with high uncertainties. Other studies suggested that ASOA
25 might have been underestimated (De Gouw and Jimenez, 2009), while some local and regional
26 studies have indicated that ASOA could be more substantial than BSOA (Aiken et al., 2009;
27 Fushimi et al., 2011). It is also thought that a substantial proportion of aerosols is formed through
28 condensation of low-volatility biogenic VOC onto existing particles of anthropogenic origin
29 (Carlton et al., 2010), thereby blurring the division between biogenic and anthropogenic SOA.
30 Several studies have indicated that anthropogenic activities could enhance the production of BSOA
31 via different mechanisms, for example by enhancing the incorporation of biogenic VOC products

1 into the condensed phase due to pre-existing organic aerosol from anthropogenic activities;
2 affecting the SOA yield by the complex effect of the anthropogenic species NO_x and nitrate (NO₃)
3 radical (a nighttime product of NO₂ and ozone) or affecting new particle formation and growth
4 where sulfuric acid has been identified as having an essential role in new particle formation (Aiken
5 et al., 2009; Carlton et al., 2010; Szidat et al., 2006; Szidat et al., 2009; Hoyle et al., 2011; Kulmala
6 et al., 2004a). Further research is thus required on the actual impact of anthropogenic emissions on
7 the formation and growth of SOA formed from biogenic VOC.

8 Several classes of VOC precursors, which are either reactive or could form oxidation products for
9 SOA formation, have been identified. Cyclic compounds are particularly important, including
10 compounds such as cycloalkanes, aromatic hydrocarbons, and terpenes. Terpenes are typically
11 oxidized by an addition mechanism, yielding oxidation products with more than two polar
12 functional groups, increasing the possibility of forming low-volatility products (Hallquist et al.,
13 2009). A number of studies have focused on the oxidation pathways of monoterpenes by hydroxyl
14 radical (OH), ozone (O₃) or nitrate radical (NO₃) forming organic acids such as pinonic and pinic
15 acid as supposed first generation oxidation products (Glasius et al., 2000; Larsen et al., 2001; Yu et
16 al., 1999; Hoffmann et al., 1997; Librando and Tringali, 2005; Surratt et al., 2008b; Calogirou et al.,
17 1999; Yasmeen et al., 2012). The first generation products are further oxidized to form more
18 oxidized products, for example pinonic acid is oxidized by the OH radical to form 3-methyl-1,2,3-
19 butanetricarboxylic acid (MBTCA) (Szmigielski et al., 2007; Müller et al., 2012). Another example
20 is the photooxidation and ozonolysis of α -pinene, one of the principal species of the monoterpene
21 class, forming terpenylic acid and diaterpenylic acid acetate (DTAA) (Claeys et al., 2009).

22 Organosulfate formation has recently been demonstrated in both laboratory studies (Surratt et al.,
23 2008b; Surratt et al., 2007a; Iinuma et al., 2007b) and ambient samples (Gómez-González et al.,
24 2008; Kristensen and Glasius, 2011; Worton et al., 2011; Gómez-González et al., 2012) as SOA
25 produced from the oxidation of biogenic VOC, such as isoprene, α -pinene, β -pinene and limonene
26 in the presence of acidic sulfate particles. In addition to representing an interesting link coupling the
27 formation of SOA from biogenic VOC precursors with the impact from anthropogenic pollutants,
28 organosulfates were also estimated to contribute up to 5-10% of the total organic mass (Tolocka and
29 Turpin, 2012) or even 30% of the total aerosol mass (Surratt et al., 2008b). Organosulfates and
30 nitrooxy organosulfates are also polar compounds, which can thus enhance the ability of aerosols to
31 act as cloud condensation nuclei (CCN) with important climate implications (Hallquist et al., 2009;
32 IPCC, 2007).

1 Here we studied the occurrence, concentrations and trends of the acidic monoterpene oxidation
2 products (including for example *cis*-pinic acid, pinonic acid, terpenylic acid, MTBCA and DTAA),
3 oxidation products from anthropogenic VOC precursors, benzoic acid and phthalic acid (typically
4 from aromatic hydrocarbons) or adipic acid and pimelic acid (from cyclic olefins), organosulfates,
5 and nitrooxy organosulfates in aerosols at two different locations in Denmark. These include an
6 urban curbside site and a semi-rural background site. The chemical analysis was performed
7 following an approach adapted from Kristensen and Glasius (2011), employing high-performance
8 liquid chromatography (HPLC) coupled with an electrospray ionization inlet (ESI) to a quadrupole
9 time-of-flight mass spectrometer (q-TOF-MS). The impacts of long-range transported SOA were
10 investigated using the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPPLIT) model
11 assisted by the modeled emissions of the major biogenic precursors including isoprene and
12 monoterpene using the global Model of Emissions of Gases and Aerosols from Nature (MEGAN)
13 and the anthropogenic emissions of sulfur dioxide (SO₂) using the long-range chemistry transport
14 model, the Danish Eulerian Hemispheric Model (DEHM), based on the European Monitoring and
15 Evaluation Program (www.EMEP.int) database. Additional support analyses were also performed,
16 including measurement of major inorganic ions to determine the aerosol acidity level, analysis of
17 local new particle formation events, local concentrations of atmospheric oxidants including ozone
18 (O₃), sulfur dioxide (SO₂) and nitrogen oxides NO_x (NO_x = NO + NO₂) and other meteorological
19 parameteres including the global radiation level, temperature, relative humidity (RH), wind
20 direction and wind speed.

21 **2 Experimental**

22 **2.1 Field campaign sites**

23 The campaign was conducted during the period May 19 - June 22, 2011 concurrently at two
24 sampling sites in Denmark (Figure 1). The sampling site HCAB (55°38' N, 12°34' E) is located in
25 central Copenhagen (~1.2 million inhabitants) on the curbside of H.C. Andersen's Boulevard with
26 considerable traffic influence (~50,000 vehicles per day) (Municipality of Copenhagen,
27 www.kk.dk). The second sampling site Risø (55°41' N, 12°05' E) is a semi-rural background site
28 located approximately 30 km west of Copenhagen and 7 km northeast of Roskilde, which houses
29 about 46,000 inhabitants. The semi-rural background area is characterized by agricultural land,
30 small villages, Roskilde Fjord located about 100 m to the west of the station, and a main road (A6)
31 located about 700 m east of the site. PM₁ samples were collected on quartz fiber filters (150 mm

1 diameter, Advantec QR-100) using Digital DHA-80 High Volume Samplers (HVS) equipped with
2 PM₁ sampling heads. The quartz fiber filters were pre-baked at 1000°C for 2 hours by the
3 manufacturer to reduce organic contamination and kept in sealed bag until use. In order to assess
4 the diurnal variation on PM₁ composition, sampling was performed in 12h daytime and nighttime
5 intervals starting every day at 06:00 am and 06:00 pm local time simultaneously at both sites. The
6 sampling flow rate was set to 23.1 m³ h⁻¹ providing a nominal sampling volume of 277.2 m³ per 12
7 h sample. While this 12 h sampling interval allowed us to sample sufficient volume of the tracer
8 compounds, the coarse time resolution complicates investigation of any short-term chemistry or
9 transport episodes. There were several gaps in the dataset with one major gap at the Risø site from
10 June 3 - 16 due to technical problems with the HVS. In total, 64 and 37 filter samples (including
11 both day and night samples) were collected at HCAB and Risø, respectively. A field blank was
12 collected in between every 12 samples or less, which showed concentrations below detection limit
13 for the organic acids, organosulfates and nitrooxy organosulfates. The field blanks were however
14 used for correction of concentrations of inorganic ions. The sample and blank filters were wrapped
15 in aluminum foil and stored in anti-static bag at - 18 °C in dark condition until analysis. **2.2**

16 **Extraction**

17 Extraction and analysis of organic acids, organosulfates and nitrooxy organosulfates followed the
18 procedure of Kristensen and Glasius (2011). Each filter sample (excluding a circle punch of 28 mm
19 in diameter used for separate analysis of ions) was extracted in 75 mL of a solution consisting of
20 90% (v/v) acetonitrile and 10% (v/v) milli-Q water by ultra-sonication for 30 minutes in a water
21 bath, where ice was regularly added to the bath to avoid heating. The extract was filtered through a
22 nylon syringe filter (0.45 µm, VWR or Q-max), evaporated to dryness using a rotary evaporator and
23 re-constituted twice in 500 µL milli-Q water containing 3% (v/v) acetonitrile and 0.1% (v/v) acetic
24 acid. Prepared samples were kept at 3-5 °C prior to analysis. Camphoric acid was used as an
25 internal standard to calculate the recovery. Field blanks were extracted and analyzed using the same
26 procedure as for the samples. It has been shown that the method could sufficiently retain more
27 volatile compounds such as pinonic acid, as well as the less volatile compounds such as pinic,
28 terpenylic and adipic acid (Kristensen and Glasius, 2011).

29 **2.3 HPLC-ESI-qTOF-MS analysis and quantification of compounds**

30 Samples were analyzed using a HPLC system (Dionex Ultimate 3000) coupled through an ESI inlet
31 to a qTOF-MS (Bruker Daltonics GmbH, Bremen, Germany). A reversed-phase HPLC separation

1 was achieved using a Waters Atlantis T3 C18 column (2.1 x 150 mm, 3 μm). The mobile phase was
2 comprised of eluent A (0.1% (v/v) acetic acid in milli-Q water) and eluent B (95% (v/v) acetonitrile
3 in milli-Q water). The injection volume was 10 μL and the flow rate of the mobile phase was 0.2
4 mL min^{-1} with the following gradient: the mobile phase was kept at 3% of eluent B in 10 min prior
5 to injection until 5 min after injection; eluent B was then increased to 20% (at 20 min), 60% (at 30
6 min) and 95% (at 35 min) where it was kept at a plateau (95%) until 45 min and increased to 100%
7 at 46 min; the gradient was kept at 100% of eluent B until 51 min and decreased to 3% (at 54 min)
8 where it was further kept at 3% for 3 min. The MS scan range was set from 50 - 1000 m/z. The
9 ESI-qTOF-MS was operated in negative ionization mode with nebulizer pressure at 3.0 bar, dry gas
10 flow 8.0 L min^{-1} , source voltage 4000 V and transfer time 50 μs . Acquired data was processed using
11 the Bruker Compass software.

12 All organic acids were quantified using authentic standards with the exception of hydroxy-pinonic
13 acid, which was quantified using pinonic acid standard. The organosulfates and nitrooxy
14 organosulfates were quantified using three different surrogate standards depending on the retention
15 time of the compound. The surrogate standards included D-mannose sulfate (molecular weight =
16 260 g mol^{-1} (MW 260)) for compounds eluting in the first 15 min, an organosulfate from β -pinene
17 (MW 250), which was synthesized in-house following Iinuma et al. (2009) for compounds eluting
18 during 15 - 40 min, and octyl sulfate (MW 210) for compounds eluting after 40 min.

19 **2.4 Extraction and analysis of ions**

20 A circle punch (28 mm diameter) of the aerosol sample filter was sonicated in 4.00 ml milli-Q water
21 for $\frac{1}{2}$ h. The extract was filtered through a polyethersulfone syringe filter (0.45 μm , PALL Life
22 Sciences IC Acrodisc) and analyzed by ion chromatography using a Metrohm 820 IC separation
23 center (with a Metrosep A Supp5-150 column for anion analysis and a Metrosep C4-150 column for
24 cation analysis) and an 819 IC conductivity detector. For anion analysis, an eluent of 3.2 mM
25 $\text{Na}_2\text{CO}_3/1.0 \text{ mM NaHCO}_3$ was used. For cation analysis, an eluent of 0.7 mM dipicolinic acid/1.7
26 mM HNO_3 was used. An auto-sampler was used where 100 μL of each sample was introduced into
27 the eluent, which was kept at a flow rate of 0.7 mL min^{-1} . The inclusion of ion analysis in this work
28 is mainly to calculate the level of acidity (H^+) from the charge balance, based on IC data of $[\text{SO}_4^{2-}]$,
29 $[\text{NO}_3^-]$ and $[\text{NH}_4^+]$ following Zhang et al. (2012b).

30 **2.5 Measurement of particle number size distributions**

1 Particle number size distributions in 15 channels from 6-700 nm were measured using a Differential
2 Mobility Particle Sizer (DMPS), which was composed of a Vienna-type Differential Mobility
3 Analyzer (DMA) (Winklmayer et al., 1991) employing re-circulating sheath air flow (Jokinen and
4 Makela, 1997) and a butanol Condensation Particle Counter (CPC) (TSI Inc. 3010, Shoreview,
5 MN). The data was corrected for reduced counting efficiency at the low size end (6-10 nm),
6 multiply charging, and losses during sampling following inversion algorithm by Wiedensohler
7 (1988). A ratio of sheath air to aerosol air of 6.8:1 was applied, with alternating up- and down-scans
8 of 150s for each size spectrum. Annual service of the CPC was performed at TSI to ensure correct
9 counting efficiency.

10 **2.6 Biogenic VOC and SO₂ emissions**

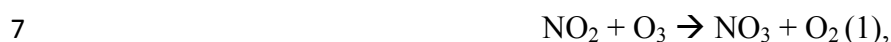
11 Emissions of biogenic VOCs are estimated by using the global model MEGAN (Guenther et al.,
12 2006; Zare et al., 2013; Zare et al., 2012). The empirical algorithm in MEGAN simulates the
13 emission rates based on relationships between ecosystem dependent emission factors and key
14 drivers that control emissions. The model takes into account the effects of temperature, radiation,
15 leaf area index, foliage age and soil moisture on the estimation of emissions. In MEGAN, plant
16 species are grouped into six plant functional types (PFTs): broadleaf tree, needle leaf evergreen tree,
17 needle leaf deciduous tree, shrub, crop and grass; and the model dataset provides the geographical
18 distribution of the standard emission factor of the six PFTs with high spatial resolution of 1 km².
19 The required meteorological inputs are provided by the MM5v3.7 model (Grell et al., 1994). The
20 initial and boundary conditions for the MM5 model are derived from the National Center for
21 Environmental Prediction (NCEP) FNL (Final Analyses) data with 6 h temporal and 1° × 1° spatial
22 resolution (<http://rda.ucar.edu/datasets/ds083.2/>).

23 Sulfur dioxide emissions were modeled using the long-range chemistry transport model, the Danish
24 Eulerian Hemispheric Model (DEHM) (Brandt et al., 2012), based on the European Monitoring and
25 Evaluation Program (www.EMEP.int) database in Europe with a resolution of 50 km × 50 km.
26 These European anthropogenic sulfur dioxide emissions are mainly released by fuel combustion
27 (www.ceip.at), dominated by coal fired power plant and international ship traffic emissions.

28 **2.7 Gas and meteorological data**

29 Meteorological data including temperature, RH and global radiation were obtained from a station
30 located on the roof-top of the H.C. Ørsted institute (HCOE, 55°42'N, 12°33'E), which represents
31 the urban background conditions of Copenhagen, Denmark. HCOE is located ~3.5 km north of
32 HCAB. O₃ and NO/NO_x were measured using photometry (Teledyne API 400, San Diego, CA,

1 USA), chemiluminescence (Teledyne API M200 A, San Diego, CA, USA) and infrared absorbance
2 (Teledyne API 300 E, San Diego, CA, USA), respectively at both field sites HCAB and Risø with
3 30 min resolution whereas measurement of SO₂ using UV fluorescence (Teledyne API M100E, San
4 Diego, CA, USA) with 30 min resolution was only obtained from HCAB. Gas and meteorological
5 data were collected within the Danish Air Quality Monitoring Program (Ellermann et al., 2012). As
6 nitrate radical is formed in the reaction:



8 the production rate of NO₃ (P_{NO₃}) was determined from the measured concentrations of NO₂ and O₃
9 using equation (2):

$$10 \quad P_{\text{NO}_3} = [\text{NO}_2][\text{O}_3]k_1 \text{ molecule cm}^{-3}\text{s}^{-1} (2)$$

11 The rate constant of reaction (1) k₁ was calculated using equation (3) with available temperature
12 data:

$$13 \quad k_1 = 1.2 \times 10^{-13} e^{-\frac{2450}{T}} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} (3) \text{ (DeMore, 1997)}$$

14 **2.8 Statistical analysis**

15 Statistical analysis was performed on the data set to analyze the intraday variability between
16 daytime and nighttime samples as well as the possible inter-site variation between the two sites
17 HCAB and Risø. As the data was not normally distributed, the U Mann-Whitney test was used
18 instead of a two-way ANOVA to assess the intraday and inter-site variability (Wilks, 2011).

19 **3 Results and Discussion**

20 **3.1 Detection and characterization of organic acids, organosulfates and nitrooxy** 21 **organosulfates**

22 During the field campaign, a total of 39 species including 12 organic acids, 18 organosulfates (OS)
23 and 9 nitrooxy organosulfates (NOS) were detected as summarized in Table 1-3. The organic acids
24 were identified using their authentic standards, with the exception of hydroxy-pinonic acid where
25 the identification was based on the characteristic mass-to-charge (m/z) fragments of the compound.
26 The organic acids included those of anthropogenic origin (i.e. benzoic acid, adipic acid, pimelic
27 acid and phthalic acid) and biogenic origin, specifically suberic acid and azelaic acid originating
28 from unsaturated fatty acids in addition to terpenylic acid, pinonic acid, pinic acid, hydroxy-pinonic

1 acid, DTAA and MTBCA originating from α/β -pinene (Table 1). Among the anthropogenic organic
2 acids, benzoic acid is a photochemical degradation product of aromatic hydrocarbons, which
3 originate from anthropogenic automobile emissions (Suh et al., 2003), while adipic acid and pimelic
4 acid have been identified as photooxidation products from the ozonolysis of cyclic olefins
5 (Hatakeyama et al., 1985; Grosjean et al., 1978; Koch et al., 2000), and phthalic acid could be the
6 oxidation product of naphthalene and other polycyclic aromatic compounds (Fine et al., 2004). The
7 biogenic dicarboxylic acids with relatively long carbon chains including suberic acid (C₈) and
8 azelaic acid (C₉) are believed to originate from the oxidation of unsaturated fatty acids (Mochida et
9 al., 2003), which are common in marine phytoplankton and leaves of terrestrial higher plants,
10 though some contribution may originate from anthropogenic meat cooking (Rogge et al., 1991) or
11 wood burning (Rogge et al., 1998). The other biogenic organic acids (terpenylic acid, pinonic acid,
12 pinic acid, hydroxy-pinonic acid and DTAA) are oxidation products of α/β -pinene (Claeys et al.,
13 2009; Ma et al., 2007), with 3-methyl-1,2,3-butanetricarboxylic acid (MBTCA) being the second
14 generation oxidation product of pinonic acid, which indicates aging by OH (Müller et al., 2012;
15 Szmigielski et al., 2007). The identified organic acids were all detected at both sites.

16 The organosulfates were identified using the characteristic fragment of m/z 97 (bisulfate anion
17 HSO₄⁻) and m/z 80 (sulfur trioxide anion SO₃⁻). The nitrooxy organosulfates were identified based
18 on a neutral loss of m/z 63 (HNO₃) in addition to the characteristic fragments of HSO₄⁻ and SO₃⁻.
19 Many organosulfates and nitrooxy organosulfates species detected in this study (Table 2-3) have
20 also been identified in previous smog chamber (Surratt et al., 2007a; Surratt et al., 2008b; Surratt et
21 al., 2010) and field studies (Gómez-González et al., 2008; Kristensen and Glasius, 2011; Gómez-
22 González et al., 2012). Most organosulfates, which were identified based on the literature probably
23 originated from isoprene (OS molecular weight (MW) 154, 156, 170, 212, 214, 216 and 334) and
24 monoterpenes including α - and β -pinene precursors (OS 248, 250, 280 and 298). Two
25 organosulfates of MW 200 were identified, of which OS 200_1 could origin from isoprene via
26 methacrylic acid epoxide (MAE) uptake onto acidified sulfate aerosol (Lin et al., 2013), while OS
27 200_2 could derive from 2-methyl-3-buten-2-ol (MBO) (Zhang et al., 2012b; Zhang et al., 2014). A
28 large share of the detected nitrooxy organosulfates could be the oxidation products from limonene
29 and/or monoterpenes (NOS 297, 311, 313, 327 and 329). The structures of several organosulfates
30 (OS 182, OS 208, and OS 210) and one nitrooxy organosulfate (NOS 331) were not identified.

31 **3.2 Concentrations**

1 The ranges of concentration levels of the detected species are shown in Table 4. In general, a higher
2 number of valid measurements (N) were available from HCAB, mostly due to the gap in the set of
3 samples collected from the semi-rural background site Risø. The total concentrations of the detected
4 organic acids averaged per sampling interval (12h) during the sampling period was $23 \pm 15 \text{ ng m}^{-3}$
5 and $19 \pm 13 \text{ ng m}^{-3}$ at HCAB and Risø, respectively, with the second generation product MBTCA
6 accounting for approximately 21% of total organic acids at HCAB and 20% at Risø. The median
7 value of MBTCA was 2.3 ng m^{-3} at HCAB and 2.0 ng m^{-3} at Risø, which are in range of the overall
8 median value of 2.7 ng m^{-3} reported by Gómez-González et al. (2012) from a study at a Belgian
9 forest site with severe urban pollution impact. Among the anthropogenic organic acids, adipic acid
10 and phthalic acid seemed to dominate at both sites, whereas azelaic acid (4.0 ng m^{-3} average) and
11 pinonic acid (2.2 ng m^{-3} average) were the dominant first generation biogenic organic acid at HCAB
12 and Risø, respectively.

13 The organosulfates were detected in most samples from both sites (Table 4). The major
14 organosulfates detected in high concentrations at both HCAB and Risø mostly originated from
15 isoprene, such as OS 154, OS 156, OS 200 and OS 212 (Surratt et al., 2007a; Surratt et al., 2008b).
16 OS 216, which was suggested to derive from the reactive uptake of isoprene epoxydiols (IEPOX) in
17 the presence of acidic sulfate aerosol (Surratt et al., 2010; Lin et al., 2012), tends to be the most
18 abundant organosulfate observed in prior laboratory studies and field studies in the southeastern US
19 (Surratt et al., 2010; Lin et al., 2012; Surratt et al., 2007b; Lin et al., 2013; Surratt et al., 2008a). In
20 this study, a moderate mean concentration of OS 216 of 5 ng m^{-3} at HCAB and 3.2 ng m^{-3} at Risø
21 was observed. It should also be noted that we used acetonitrile as extraction solvent instead of
22 methanol as used by Surratt et al. (2007b, 2008, 2010) and Lin et al. (2012, 2013), which could
23 have affected the extraction yield of OS 216 in the present work. The total organosulfate
24 concentration averaged per sampling interval (12h) during the sampling period was $60 \pm 45 \text{ ng m}^{-3}$
25 at HCAB, whereas at Risø a slightly lower total concentration value of $47 \pm 31 \text{ ng m}^{-3}$ was obtained.
26 Nitrooxy organosulfates were detected in lower concentrations compared to organosulfates, with a
27 total concentration of $4 \pm 1 \text{ ng m}^{-3}$ at HCAB and $3 \pm 1 \text{ ng m}^{-3}$ at Risø. NOS 297, which was possibly
28 an oxidation product from limonene (Surratt et al., 2008b) was the major nitrooxy organosulfate
29 found at both sites.

30 The concentration levels of the detected compounds were compared with an earlier study performed
31 at a forest site in Denmark by Kristensen and Glasius (2011) during spring 2008, in which fewer
32 species (six organic acids, five organosulfate, five nitrooxy organosulfates) were included. The

1 comparison was thus only targeted at similar compounds. In general, the concentrations of most
2 organic acids were higher in the present study, except *cis*-pinic acid and pinonic acid, which were
3 more abundant at the forest site. The concentration levels of the organosulfates and nitrooxy
4 organosulfates in this study were many-fold higher compared to the previous study, especially with
5 regards to the organosulfate species. This, in addition to the differences among the sites and years
6 could be caused by differences in quantification standards, as Kristensen and Glasius (2011) used
7 camphor sulfonic acid as a surrogate standard, as opposed to the three different standards including
8 D-mannose sulfate, an in-house synthesized organosulfate from β -pinene and octyl sulfate used in
9 this study (as described above). This highlights the analytical challenge of quantifying
10 organosulfates and comparing concentrations among different studies.

11 The total concentration of organic acids and organosulfates are significantly higher in our samples
12 compared to the Arctic, where a total concentration of $3 \pm 1 \text{ ng m}^{-3}$ for organic acids and $12 \pm 2 \text{ ng}$
13 m^{-3} for organosulfates (quantified using the same organosulfate from β -pinene used in the present
14 study) were measured at Zeppelin Mountain, Svalbard during summer of 2008 (Hansen et al.,
15 2014). However, during late winter and early spring, total concentration levels up to 70 ng m^{-3} and
16 over 45 ng m^{-3} were detected for all organic acid, organosulfate and nitrooxy organosulfate
17 compounds respectively at Station Nord, northeast Greenland and at Zeppelin (Hansen et al., 2014).
18 These values are in range with our concentration values. It must be noted that a total of 11 organic
19 acids, 12 organosulfates and one nitrooxy organosulfates were detected by Hansen et al. (2014).

20 **3.3 Inter-site, day/night variability and inter-species correlation**

21 The non-parametric U Mann-Whitney test was performed on the data sets to analyze the inter-site
22 variability between HCAB and Risø during the overlapping period (May 19 - June 1 and June 17 -
23 22) where data from both sites were available. The 90% confidence interval was used for evaluation
24 of the statistical significance to account for the possible field sampling, filter extraction and
25 quantification uncertainties. The test results showed that 14 parameters out of the 41 tested
26 parameters were statistically different between the two sites at 90% confidence interval ($p < 0.1$)
27 (Supplementary Table 4). These 14 parameters included six organic acids (benzoic acid, pimelic
28 acid, phthalic acid, suberic acid, pinonic acid, and azelaic acid), the concentrations of total first
29 generation organic acids, five organosulfates (OS 182, OS 200, OS 210, OS 248 and OS 298), one
30 nitrooxy organosulfate (NOS 297) and the concentration of total nitrooxy organosulfates. The
31 species with statistically higher concentrations were mostly from HCAB (12 out of 14) with the

1 exception of benzoic acid and pinonic acid, which showed significantly higher concentrations at
2 Risø. While a higher occurrence of benzoic acid at the semi-rural background site Risø was
3 unexpected, it seems that the urban curbside site HCAB was associated with enhanced
4 concentrations of both biogenic (suberic acid and azelaic acid) and anthropogenic (pimelic acid and
5 phthalic acid) compounds in addition to the anthropogenic-biogenic coupling species (OS 182, OS
6 200, OS 210, OS 248, OS 298 and NOS 297), which probably indicated some local anthropogenic
7 enhancement impacts on the concentration levels of the compounds. In fact, the total concentrations
8 of organic acids and nitrooxy organosulfates were also found at significantly higher concentrations
9 at HCAB. In contrast, the second generation product of pinonic acid, MBTCA, which has been
10 regarded as an indicator of photochemical aging of pinonic acid by OH (Müller et al., 2012) was not
11 statistically different between the two sites. The fact of a statistically higher concentration level of
12 the pinonic acid precursor at Risø and the indifferent concentration of its product MBTCA at the
13 sites probably suggest that the degree of photochemical aging was not directly governed by the
14 concentrations of the precursor at the semi-rural background site Risø.

15 The variability between daytime and nighttime samples at each site during the whole field study
16 was also analyzed using a non-parametric U Mann-Whitney test at 90% confidence interval. The
17 test results showed statistically indistinguishable daytime and nighttime concentrations for most
18 species (Supplementary Table 1-2), with the exception of higher daytime concentrations of phthalic
19 acid ($p = 0.0385$), OS 210 ($p = 0.0622$) and NOS 343 ($p = 0.0825$) at HCAB and higher nighttime
20 concentrations of NOS 297 at both sites ($p = 0.0011$). Phthalic acid, OS 210 and NOS 297 were
21 associated with high numbers of valid measured daytime or nighttime concentrations ($N \geq 30$)
22 whereas NOS 343 was associated with a lower number of valid N ($N_{\text{day}} = 10$; $N_{\text{night}} = 13$).

23 Phthalic acid is widely used in plasticizer substrates and could be primarily emitted to the
24 atmosphere from off-gassing of plasticizers from plastics (Fraser et al., 2003), though previous
25 studies have also suggested that it correlated with SOA in ambient samples (Fraser et al., 2003; Fine
26 et al., 2004; Schauer et al., 2002). A recent study of the photooxidation of naphthalene has also
27 indicated the link between phthalic acid formation and photochemistry, while pointing towards
28 negligible primary emissions of the compound based on laboratory and field measurements
29 (Kleindienst et al., 2012). The findings support our observation of higher daytime concentration of
30 phthalic acid at HCAB, though phthalic acid was not necessarily formed directly at the site.
31 Meanwhile the indifferent daytime and nighttime concentrations of phthalic acid at Risø were
32 probably governed by a generally lower concentration level of the species at the site

1 (Supplementary Table 4). The other compound with significantly higher daytime concentrations at
2 HCAB was OS 210, which has recently been reported in aerosols in the Arctic (Hansen et al.,
3 2014).

4 The compound NOS 297 (m/z fragments 296, 233, 177, 163 and 97) showed significantly higher
5 nighttime concentration at both sites, though its concentrations were statistically higher at HCAB
6 compared to Risø (Supplementary Table 4). This species was previously detected in ambient
7 aerosols (Gao et al., 2006; Reemtsma et al., 2006; Kristensen and Glasius, 2011) and has been
8 suggested as an oxidation product of limonene-like monoterpene precursors (Surratt et al., 2008b).
9 In this study, NOS 297 was identified as the major nitrooxy organosulfates (Table 4), accounting
10 for 36% and 34% of the total concentrations of nitrooxy organosulfates detected at HCAB and Risø,
11 respectively. The higher nighttime occurrence of NOS 297 could indicate the impacts of local or
12 regional nighttime chemistry on the concentration level of the compound. In fact, the formation of
13 nitrooxy organosulfates has been suggested to involve nighttime nitrate (NO₃) radical chemistry
14 (Iinuma et al., 2007a; Iinuma et al., 2007b; Surratt et al., 2008b). In contrast, NOS 343 showed
15 significantly higher daytime concentrations at HCAB, though the statistical calculation was
16 associated with a lower number of measurements, as mentioned above. Too few measurements of
17 NOS 343 were available at Risø to compare its daytime and nighttime concentrations. Overall,
18 significantly higher nighttime concentrations were observed for total nitrooxy organosulfates at
19 both sites (Supplementary Table 1-2), which was attributed to the predominant contribution from
20 the nighttime-dominant species NOS 297. As a majority of the detected species did not show any
21 immediate evidence of local site impacts on their concentration levels according to the U Mann-
22 Whitney test results, the relations among the compounds were furthermore investigated using R²
23 correlation coefficients to unveil any potentially driving mechanisms or factors influencing their
24 formation. The correlation coefficients were calculated for each pair of species at each site and
25 subsequently averaged for both sites. The averaged R² correlation coefficients are presented in
26 Figure 2, with each organic acid, organosulfate and nitrooxy organosulfate compound (lines)
27 correlating with other species (x-axis). Relative standard deviations (RSD, %) for the corresponding
28 averaged R² correlation coefficient are shown in Supplementary Table 1. Only 36 out of the total 38
29 species were used to calculate the R² correlation coefficient values as those with very few
30 measurement values were excluded. The results in Figure 2 were divided into panels of organic
31 acids (top), organosulfates (middle) and nitrooxy organosulfates (bottom).

1 Figure 2 shows a clear correlating group with high R^2 correlation coefficient values (> 0.5) among
2 the species in the group, which is comprised of certain organic acids, organosulfates and nitrooxy
3 organosulfates. The correlating group included three organic acids, namely terpenylic acid,
4 MBTCA and DTAA (dark blue), a vast number of 12 organosulfates (red) and four nitrooxy
5 organosulfates (green). These organosulfates and nitrooxy organosulfates among others included
6 species closely related in molecular weight, such as the groups of OS (154, 156), OS (212, 214,
7 216), OS (252, 254) and NOS (327, 329, 331). The organic acids pinonic and pinic acid (light blue)
8 and OS 248 (purple) seemed to only partly follow the correlation pattern as the degree of correlation
9 was somewhat lower, especially with regards to the correlation between these species and
10 organosulfates.

11 The degree of correlation (calculated as the R^2 values) among the mentioned specific compounds
12 was persistent at both study sites, as evident by the generally lower deviation with small RSD
13 values (%) associated with averaging the R-squared values from both sites as opposed to the
14 uncorrelated species (Supplementary Table 3). The organosulfates correlated consistently well with
15 the organic acids and the other organosulfates within the correlating group, with RSD ranging from
16 typically a few to over 20% between sites. Correlation coefficients between the nitrooxy
17 organosulfates and the other species showed larger discrepancies between the two sites, with RSD
18 ranging from a few percent (NOS 313) to considerably higher percentages (NOS 327, 329 and 331),
19 mostly due to a higher correlation level of these species at Risø compared to HCAB. Concentrations
20 of pinonic acid, pinic acid and phthalic acid, which only partly followed the correlation pattern
21 showed larger deviations between the two sites. In contrast, OS 248 which also partly followed the
22 pattern (with slightly lower R^2 coefficient ~ 0.5) showed a highly consistent correlating tendency at
23 both sites with low RSD values. The consistent correlation pattern among the group of organic
24 acids, organosulfates and nitrooxy organosulfates of different precursors detected at both sites
25 probably indicate a common source for the species at both sites.

26 **3.4 Regional impacts**

27 Figure 3 shows the temporal variation of the total concentrations of organic acids, organosulfates,
28 nitrooxy organosulfates and PM_{10} detected at HCAB and Risø during the field study. It is apparent
29 that all three classes of compounds showed similar temporal patterns, and the two sites do not differ
30 substantially, which were attributed to the large contribution of compounds belonging to the
31 correlating group affecting both sites. The observations strongly indicated that the major sources or
32 chemistry governing the total concentration levels of the compounds must occur at a spatial location

1 affecting both the urban curbside and semi-rural background sites, which represent quite different
2 environments (Figure 1). Elevated total concentrations of organic acids, organosulfates and nitrooxy
3 organosulfates seemed to occur at both sites during the same daytime or nighttime interval,
4 exemplified by the daytime samples on May 22, May 31 and June 18 or the nighttime sample on
5 May 23. Several other concentration maxima occurring on June 6 (daytime) and June 8 (nighttime)
6 were observed at HCAB (data from Risø was unavailable for these days). Organosulfate 200_2
7 (from MBO), which was mostly found at levels below the detection limit, was detected in
8 considerably higher concentrations during these episodes of concentration maxima (for example in
9 May 31, June 18 (daytime) and May 23 (nighttime) samples).

10 The correlation between the total concentration level of organosulfates and the concentration of
11 SO_4^{2-} ion measured by IC was also examined. A relatively high correlation coefficient value ($R^2 =$
12 0.6) was obtained between the total organosulfates and sulfate at the HCAB site with the episodes
13 of highest concentrations of organosulfates coinciding with peak concentrations of the SO_4^{2-} ion
14 (Figure 3), which is in line with previous observations of aged continental aerosols over the
15 southeast Pacific Ocean (Hawkins et al., 2010). The correlation between organosulfates and
16 inorganic sulfate also agrees with previously proposed formation mechanisms for organosulfates
17 including reactive uptake of epoxides on the surface of acidic sulfate aerosols (Iinuma et al., 2009;
18 Minerath and Elrod, 2009; Surratt et al., 2010; Lin et al., 2013; Worton et al., 2013) and sulfate
19 radical-initiated formation in the bulk phase (Nozière et al., 2010). However, while this observation
20 may support related formation processes of the organosulfates and the sulfate ion, the resemblance
21 in concentration trend among sulfate, organosulfates, organic acids, and nitrooxy organosulfates
22 may also indicate that organic acids, organosulfates and nitrooxy organosulfates could also partition
23 into existing sulfate aerosol. It should be noted that the correlation was only visible at the urban
24 curbside site HCAB, while such correlation was not found at Risø ($R^2 = 0.3$).

25 In general, the production rate of NO_3 radical was one order of magnitude higher at the urban site
26 HCAB compared to the semi-rural background Risø, which was also observed in an earlier study in
27 the region and attributed to higher emissions of NO_2 in urban condition (Ljungstrom and Hallquist,
28 1996). Although the formation of nitrooxy organosulfates could involve nighttime NO_3 radical
29 chemistry as discussed above, no direct correlation was found between the nighttime total
30 concentration level of nitrooxy organosulfates and the corresponding production rate of NO_3 radical
31 at HCAB ($R^2 = 0.1$). Meanwhile, a good correlation ($R^2 = 0.6$) of nighttime P_{NO_3} and nighttime total
32 concentration of nitrooxy organosulfates was found at Risø (Figure 3). The difference could be

1 attributed to the continuous emissions of NO from automobiles in the city centre, which reacts with
2 NO₃ radical and complicates the observed trend of its production rate.



4 In addition, the observed concentrations of nitrooxy organosulfates could also be affected by both
5 local and long-range transported impacts. The extent and impact of long-range transport, the impact
6 of the interplay between O₃, NO₂ and the production rate of NO₃ radical on the possible local
7 species will be discussed in more detail below.

8 In order to investigate the regional impacts on the concentration levels of the detected compounds,
9 5-days air mass back trajectories were calculated using the Hybrid Single Particle Lagrangian
10 Integrated Trajectory (HYSPLIT) model for every 12h sample (Draxier and Hess, 1998). As HCAB
11 and Risø are relatively close with a distance of ~30 km, HYSPLIT calculations were only
12 performed for the HCAB site. The results revealed a dominating westerly origin of air masses
13 during most of the campaign period. However a clear dominance of the south-easterly air masses
14 was found to be associated with the daytime or nighttime episodes of elevated concentrations,
15 including the daytime samples on May 22, 26 and 31 and June 6 and 18 and the nighttime sample
16 on June 8. The travelling path of the south-easterly air masses probably involved the region around
17 the borders between Germany, Poland and Czech Republic (Figure 1, 4). It must be noted that the
18 air masses following the south-easterly direction could have different origins (3-5 days backwards)
19 but all passed the south-easterly region 1-2 days immediately before arrival at the sampling site. An
20 example of the impacts of air masses arriving from the south-easterly direction is shown for three
21 consecutive days on May 30th (nighttime), May 31 (daytime with elevated concentrations) and June
22 1 (daytime) where data from both HCAB and Risø were available (Figure 4). To support the
23 interpretation of the HYSPLIT calculations, emissions of isoprene and monoterpenes (daily)
24 calculated using the MEGAN model and SO₂ (monthly) calculated using the DEHM model, are also
25 shown in Figure 4 to illustrate the magnitude of biogenic and anthropogenic emissions in the
26 region.

27 Figure 4 shows a sudden change of air mass origin from westerly (May 30) to south-easterly (May
28 31) back to westerly (June 1), which was coupled with a major episode on May 31 with elevated
29 concentrations of the organic acids, organosulfates and nitrooxy organosulfates (Figure 3). The
30 modeled isoprene and monoterpene daily emissions did not vary noticeably over the corresponding
31 days (Figure 4), while emissions from the westerly direction affecting May 30 and June 1 seemed

1 comparable to emissions from the southeasterly direction affecting May 31. This indicated that
2 VOC emissions were probably not the limiting factor in the formation of the investigated
3 compounds.

4 Figure 4 also shows a map of average SO₂ emission over the two months May - June 2011. Though
5 the SO₂ emission map could not be used to evaluate the daily change in anthropogenic emissions to
6 further investigate the observed elevated concentrations on May 31, it however indicated strong SO₂
7 emissions from hotspots in the region that were passed over by the south-easterly air mass shown in
8 the trajectories. This could probably influence the formation of the organosulfates and nitrooxy
9 organosulfates during long-range transport to the sites. The accompanying organic acids, which
10 were also found at high concentrations correlating to the organosulfates and nitrooxy
11 organosulfates, could also be long-range transported to the sampling sites.

12 It is difficult to determine the exact spatial location of the major sources or chemistry governing the
13 total concentrations of the compounds at the two sites, which is partly due to the coarse resolution
14 of the SO₂ emission map. However, it is unlikely that the major source location would be in the
15 local proximity of the two sites due to the highly different local conditions between the urban
16 curbside site HCAB and the semi-rural background site Risø (Figure 1). Such a local source, if
17 present, would be subject to immediate different local background conditions and also varying local
18 wind conditions, while in contrast, a highly similar temporal variation pattern of total observed
19 concentrations was found at the sites during the whole campaign period (Figure 3). Furthermore,
20 any possible point source located southeasterly from HCAB and Risø, which could affect both
21 sampling sites would unlikely be in Denmark, as both sites are located only 10 - 20 km from Baltic
22 sea in the southeasterly direction (Figure 1). The source region therefore is possibly located at the
23 broader regional scale across the Baltic sea extending to the southerly neighboring countries.

24 Particle mass concentrations were calculated based on integration of the calculated particle mass
25 size distributions (from DMPS data), which were further based on measured particle number size
26 distributions assuming particle sphericity, using a SOA particle density of 1.7 g cm⁻³ (Stock et al.,
27 2011) (Figure 3). This density value is close to the bulk densities of ammonium sulfate and nitrate,
28 which are expected to be the major components of aged accumulation mode particles. The detected
29 and quantified organosulfates and nitrooxy organosulfates contributed to 0.1 - 1.0 % (approximately
30 0.5 % on average) of PM₁ mass at HCAB and 0.4 - 1.5 % (approximately 0.8 % on average) of PM₁
31 mass at Risø. The values are considerably lower than previous estimates, including 30% (Surratt et

1 al., 2008b) or 5-10% contribution of organosulfates to total organic mass (Tolocka and Turpin,
2 2012). PM₁ concentrations at HCAB were in the lower range during the days under the influence of
3 southeasterly air mass (May 22, May 26, May 31, June 18 (daytime), with the exception of June 6
4 (daytime). The period of high PM₁ concentrations at HCAB (June 6 - 8) seemed to link to other
5 local conditions, including episodes of elevated concentrations of SO₂ and NO_x (Figure 5 and
6 Supplementary Figure 1). The local impacts are further discussed in the section below.

7 **3.5 Local impacts**

8 The possible local anthropogenic impacts were investigated by examining the species of
9 significantly higher concentrations at HCAB, which did not belong to the correlating group,
10 including OS 182 (m/z fragments 181, 97), OS 210 (m/z fragments 209, 153, 137, 97 and 79) and
11 NOS 297 (m/z fragments 233, 177, 163 and 97). As displayed in Figure 5, the patterns of changes
12 in OS 182 and OS 210 concentrations in May were considerably similar, with a few episodes of
13 high daytime concentrations especially during the period May 20 - 25 with a lower concentration on
14 May 22 (daytime). Such observation stands in contrast to the trend of the total organosulfate
15 concentrations during the same period, where a generally lower concentration level was observed
16 except from elevated daytime concentrations on May 22 (Figure 3). Assuming that the trend of
17 variations the total detected species in Figure 3 was mostly governed by long-range transported
18 compounds belonging to the highly correlated group, the trend of OS 182 and OS 210, which did
19 not belong to the correlated group, could probably reflect some local impact on the concentration
20 levels of these species. However, while OS 182 and OS 210 seemed to correlate well in May,
21 concentrations of the species deviated towards the end of the field study (Figure 5), which could be
22 partly attributed to additional input from long-range transport to the site. For example, elevated
23 daytime concentration of OS 182 was observed on June 16, which is one of the characteristic days
24 of southeasterly arriving air masses associated with higher total concentrations of the detected
25 species (Figure 3).

26 Local impacts were studied to explain the significantly higher concentrations of OS 182, OS 210
27 and NOS 297 at the urban curbside site HCAB. Temporal variation of gases (NO, NO_x and SO₂),
28 aerosol acidity level (calculated from charge balance using [SO₄²⁻], [NO₃⁻], and [NH₄⁺] following
29 Zhang et al. (2012b)) at HCAB (Figure 5), and meteorological conditions including RH (Figure 5),
30 temperature (Temp) and global radiation (GR) (Supplementary Figure 1) at the urban background
31 site HCOE, which was located in close proximity to HCAB were analyzed. As can be seen from
32 Figure 5, the period May 20 - 25 was accompanied by a generally higher level of aerosol acidity

1 which decreased on May 22 (daytime). Such variation of the aerosol acidity level perfectly matched
2 the trend of the varying concentrations of OS 210 and OS 182 during the same period (Figure 5). In
3 fact, the aerosol acidity level has been suggested as an essential parameter in the formation of
4 organosulfates (Zhang et al., 2012b). At the same time, the concentration level of the organosulfates
5 might rather be affected by an inter-play between the aerosol acidity level and other factors, such as
6 the global radiation and temperature. For example the May 20 and 21 (daytime) samples showed
7 high concentrations of OS 182 and OS 210, which was comparable to the concentrations on May 24
8 or 25 (daytime) samples (Figure 5), whereas the aerosol acidity level on May 20 or 21 was
9 considerably lower on May 20 and 21 (daytime) compared to the other days (Figure 6). May 20 - 21
10 were however accompanied with a higher level of global radiation, especially on May 21 (579 W m^{-2})
11 compared to other latter days May 24-25 (daytime) (Supplementary Figure 1). While a higher
12 level of global radiation could probably enhance the oxidation of SO_2 by the OH radical and thereby
13 resulted in higher concentrations of SO_4^{2-} as observed with the May 20 - 21 (daytime) samples,
14 NH_4^+ was also detected at higher concentrations with these samples, resulting in an overall lower
15 aerosol acidity calculated based on ion charge balance on the days. In addition, the higher
16 temperature observed on May 20 - 21 compared to May 24 - 25 may also have contributed to
17 enhance the formation of supposed local organosulfates in the former samples, by means of
18 enhancing biogenic VOC emissions, or photochemistry in general. In fact, a considerably higher
19 level of O_3 oxidant was also found during the former days (May 20 - 21) compared to the latter days
20 (May 24 - 25), which would positively enhance the formation of SOA.

21 Higher daytime concentrations of OS 210 were also observed on June 7 and 8, whereas the aerosol
22 acidity reached its highest level of 46 nmol m^{-3} on June 7 (nighttime) with interestingly almost
23 neutral conditions on the adjacent June 7 (daytime) and June 8 (daytime) (Figure 5). At the same
24 time, during these two daytime intervals, higher average concentrations of SO_2 were also observed.
25 At the same time, concentrations of NO and NO_x in higher resolution (30 min) were also
26 considerably elevated (Supplementary Figure 1). Meanwhile, the O_3 level was in the lowest
27 concentration range presumably due to the lower level of sunlight in addition to titration by NO_x
28 (figure not shown). A close examination revealed that the NO and NO_x were highest during the
29 early morning and late afternoon hours which are attributed to traffic emissions, though during the
30 day course both NO and NO_x remained at a higher level compared to the other days during the
31 campaign period. This observation could be partially attributed to lower wind speed during daytime
32 on June 7 (2.0 m s^{-1}) and on June 8 (2.9 m s^{-1}), which was considerably lower than the average wind

1 speed of 4.1 m s^{-1} for the whole campaign period. It must also be noted that this period was coupled
2 with the highest RH range during the campaign period (Supplementary Figure 2), which resulted
3 from the corresponding very low global radiation during the days. The observations indicated an
4 inter-play between the anthropogenic gases, especially NO, NO_x and probably aqueous aerosol
5 chemistry, leading to the occurrence of higher concentrations of OS 210 in these samples. A
6 previous study by Dommen et al. (2006) on the effect of RH under high NO_x conditions indicated
7 that more volatile SOA were observed at higher RH rather than at lower RH. Additionally, SOA
8 formation including carboxylic acids and organosulfur compounds in cloud and fog droplets have
9 been hypothesized as plausible by Blando and Turpin (2000) while a recent smog chamber study by
10 Zhang et al. (2012a) suggested that elevated RH values could mediate the formation of
11 organosulfates, thereby emphasizing the role of wet sulfate aerosols in forming organosulfates in
12 ambient aerosols. At the same time, though aerosol acidity has been found to enhance isoprene
13 SOA formation under low initial NO conditions (Surratt et al., 2010; Surratt et al., 2007c; Jaoui et
14 al., 2010), seed aerosol acidity showed a negligible effect on SOA formation from isoprene under
15 high initial NO conditions (Surratt et al., 2006), which implied the less important role of seed
16 aerosol acidity on the formation of OS 210 on June 7 and 8 (daytime) where both NO and RH were
17 high.

18 Further investigation of the meteorological conditions during the period where data from both sites
19 were available (May 19 - June 1 and June 17 - June 22, see Supplementary Figure 2) revealed a
20 possible coupling between higher temperature and global radiation (508 Wm^{-2}) with the occurrence
21 of concentration maxima observed on days with the highest temperature May 31 (daytime) and June
22 6 (daytime) (Figure 3). However, information was lacking on the spatial scale of high temperature
23 and global radiation episodes to fully justify if concentration maxima were enhanced by high
24 temperature and global radiation at source or at the sampling sites.

25 The third species used to study the local impacts was NOS 297, which showed significantly higher
26 nighttime concentration as discussed above. A recent work by Rollins et al. (2013) has also
27 suggested a correlation between nitrooxy organosulfates, including NOS 297 and total particulate
28 organic nitrate (RONO₂), which results from oxidation of organic compounds in high NO_x
29 environment and NO₃ radical chemistry. Although some possible long-range transport contribution
30 to the observed concentration of NOS 297 at HCAB could not be completely eliminated, as shown
31 by its elevated concentration seen during peak concentration days May 31 and June 16, elevated
32 concentrations of NOS 297 were observed predominantly at nighttime on May 19, 20, 22, 26 and

1 June 6, 8 and 10. To examine the local impacts on NOS 297, only nighttime production rate of NO₃
2 radical and nighttime RH are shown (Figure 5). The RH condition during the nights of high
3 concentrations of NOS 297 covered both the lower RH range (May 19, May 20 and June 10) and
4 the higher RH range (May 22, June 6 and June 8) of the whole campaign. Very high production rate
5 of NO₃ radical (12.8 and 9.7 x 10⁷ molecule cm⁻³ s⁻¹, respectively) was observed during the nights
6 of highest RH, June 6 and 8, whereas during the in-between night June 7, when RH was still in the
7 high range, lower P_{NO₃} was observed, which was coupled with considerably lower concentration of
8 NOS 297. In contrast, lower P_{NO₃} was also observed during other nights of higher NOS 297
9 concentrations with lower RH condition (May 19, 20 and June 10). However, a lower RH range
10 could also reduce the available condensation sink for particle precursors and thus facilitate the
11 formation of NOS 297 by nighttime NO₃ radical chemistry. It is thus unlikely if any single
12 parameter such as RH or P_{NO₃} could be a prerequisite for the occurrence of elevated concentration
13 of NOS 297 in this study. In addition, investigation of air mass back trajectories using HYSPLIT
14 also revealed that the May 22, 26 and June 6 nighttime were affected by air masses arriving from
15 the south-south-easterly direction whereas on June 8 air masses arrived from easterly direction.
16 Such observations suggest that the detected nighttime concentrations of NOS 297 could be
17 governed by both the local formation and the degree of long-range transport of the compound to the
18 site. As a result, no direct correlation between nighttime concentrations of NOS 297 and P_{NO₃} was
19 observed (R² = 0.2). On the other hand, it should be noted that during this period of the year, the
20 duration of the night is relatively short in Denmark, therefore calculation of nighttime
21 concentrations would couple with higher uncertainty. Nevertheless, it seems probable that
22 occurrence of NOS 297 was favored by nighttime chemistry, regardless of whether the species were
23 formed in-situ or long-range transported to the site.

24 Local new particle formation (NPF) events were also examined using the DMPS data following Dal
25 Maso et al. (2005) to investigate whether the local formation of organosulfates could be associated
26 with particle nucleation at the sites. A limited number of NPF events were found occurring on the
27 same days at the semi-rural background site Risø, the urban background site HCOE and the urban
28 curbside site HCAB, though HCAB is heavily influenced by traffic emissions resulting in generally
29 short nucleation bursts rather than allowing a classic “banana” formation and growth. The NPFs
30 however seemed to occur on the days with low concentrations of most organic acid, organosulfate
31 and nitrooxy organosulfate species, including the possible local compounds OS 182 and OS 210,

1 which agree with previous indications that NPF may occur more frequently in cleaner environments
2 (Lyubovtseva et al., 2005; Kulmala et al., 2004b). As the occurrence of higher concentrations of the
3 detected organic acids, organosulfates and nitrooxy organosulfates did not correspond to NPF
4 events, which were coupled with elevated occurrence of smaller-sized particles; it is likely that the
5 organic acids, organosulfates and nitrooxy organosulfates belong to the accumulation mode. At the
6 same time, any correlation to NPF should be investigated with size-segregated samples.

7 **4 Conclusion**

8 A field study was conducted from May 19 to June 22, 2011 at two sites in Denmark, including the
9 urban curbside site HCAB and the semi-rural background site Risø to investigate the anthropogenic
10 impacts on the formation of biogenic SOA via analysis of organosulfates and related oxidation
11 products using HPLC-qTOF-MS. A substantial range of anthropogenic and biogenic organic acids,
12 organosulfates and nitrooxy organosulfates were detected, identified and quantified. Isoprene
13 oxidation products comprised a large fraction of the detected organosulfates whereas a majority of
14 the detected nitrooxy organosulfates may originate from limonene and/or monoterpenes. The U
15 Mann-Whitney statistical test showed significantly higher concentrations of specific biogenic and
16 anthropogenic species at HCAB, whereas an intraday statistical test only revealed significantly
17 higher daytime concentrations of phthalic acid and OS 210 and significantly higher nighttime
18 concentrations of NOS 297. Many detected compounds including a group of organic acids,
19 organosulfates and nitrooxy organosulfates of various precursors were found belonging to a
20 common highly correlated group, consistently affecting their concentration levels at both sites,
21 suggesting a common source region or similarities in formation processes. The analysis of
22 HYSPLIT back trajectories together with isoprene and monoterpene emissions calculated by the
23 MEGAN model and SO₂ emission based on the EMEP monitoring database indicated that south-
24 easterly air mass arriving from across the Baltic sea could relate to the corresponding episodes of
25 the elevated total concentrations at the sites, implying the importance of the regional impacts on the
26 occurrence of the compounds.

27 The local impacts were also investigated for several selected species which showed significantly
28 higher concentrations at HCAB compared to Risø and did not belong to the correlating group,
29 including OS 182, OS 210 and NOS 297. The results showed that the local factors including aerosol
30 acidity, RH, NO and probably some wet aerosol chemistry could possibly lead to a higher
31 occurrence of the organosulfates, whereas the major nitrooxy organosulfate NOS 297 could be

1 favoured by nighttime chemistry, in addition to temperature and global radiation, which could also
2 enhance the concentrations of the long-range transported species at the sites. As only very few
3 compounds were deemed as affected by the local conditions, the analysis of the local impacts was
4 restricted to a modest number of compounds. In general, the regional impacts seemed to
5 considerably exceed the local impacts. It was estimated that organosulfates and nitrooxy
6 organosulfates contributed to approximately 0.5% and 0.8% of PM₁ mass, respectively at HCAB
7 and Risø.

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10 providing the meteorological and gas data from The Danish Air Quality Monitoring Program.

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1 **List of Figures**

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4 centre of Copenhagen close nearby a busy street. Risø is located approximately 30 km west of
5 HCAB.

6 Figure 2. Correlations between organic acids, organosulfates and nitrooxy organosulfates with each
7 other detected species in each sample throughout the sampling period (excluding those with too few
8 measurements), expressed as correlation coefficient (R^2). The results show R^2 coefficient averaged
9 from the two sites. High R^2 values (> 0.5) were found among a specific group of species (yellow
10 shade), suggesting common sources. A few other species including pinonic acid, pinic acid, phthalic
11 acid, and OS 248 also partly followed the correlation pattern.

12 Figure 3. Temporal trends of total concentrations of organic acids, organosulfates, nitrooxy
13 organosulfates and PM_{10} at the urban curbside site HCAB (solid line) and the semi-rural background
14 site Risø (broken line). The figure shows a strong correlation between the two sites and among the
15 three categories of compounds examined. Sulfate (HCAB) and production rate of NO_3 radical
16 (Risø) are also shown in the organosulfate and nitrooxy organosulfate panels, respectively.

17 Figure 4. A sudden change in air mass back trajectories on May 31 to southeasterly direction was
18 coupled with elevated concentrations of organosulfates, nitrooxy organosulfates, and organic acids
19 at both HCAB and Risø. Modeled daily isoprene and monoterpene emissions did not vary
20 significantly over the three days. The SO_2 emission map (May-June average) showed a higher
21 abundance of emission hotspots in southeasterly direction (observed on May 31) compared to the
22 westerly direction (observed on May 30 and June 1). HCAB is marked at the bottom tip of the black
23 triangle.

24 Figure 5. Time profiles of selected species of significantly higher concentration levels at HCAB,
25 including OS 182, OS 210 and NOS 297. Concentrations of OS 182 and OS 210 are shown against
26 level of SO_2 and acidity at HCAB, while concentration trend of NOS 297 is shown against
27 nighttime production rate of NO_3 radical and nighttime relative humidity at HCAB.

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4 Table 2. Detected organosulfates (OS) during the campaign.

5 Table 3. Detected nitrooxy organosulfates (NOS) during the campaign. Table 4. Concentration
6 range (ng m^{-3}) of the detected species at HCAB and Risø, reported as mean, standard deviation
7 (stdev.), median, max and the number of detected samples (N) at each site.

8 **Supplementary Information**

9 Supplementary Table 1. Statistical results of U Mann-Whitney test performed on daytime and
10 nighttime concentrations of the organic acids, organosulfates and nitrooxy organosulfate detected at
11 HCAB. Statistically different parameters (with p-level < 0.1) are underlined. Parameters with two
12 few measurements ($N < 5$) were excluded from the test.

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15 nighttime concentrations of the organic acids, organosulfates and nitrooxy organosulfate detected at
16 Risø. Statistically different parameters (with p-level < 0.1) are underlined. Parameters with two few
17 measurements ($N < 5$) were excluded from the test.

18 Supplementary Table 3. Relative standard deviations (RSD, %), which were associated with
19 averaging R-square correlation coefficients (between pairs of compounds) at the two sites, were
20 calculated. The species belonging or partly belonging to a common correlating group are marked in
21 blue shade with the corresponding RSD in red (those only partly belonging to the group
22 underlined). RSD values seemed low for the correlating organic acids and OS while larger for the
23 NOS, mostly due to higher correlation coefficients at Risø. The partly-correlating species pinic,
24 pinonic acid and phthalic acid showed relatively larger RSD compared to the other correlating
25 compounds, while another partly-correlating species (OS 248) showed a consistent partly
26 correlating degree at both sites as evident by small RSD values.

27 Supplementary Table 4. Statistical results of U Mann-Whitney test performed on the organic acids,
28 organosulfates and nitrooxy organosulfate detected at the two sampling sites H.C. Andersen
29 Boulevard (HCAB) and Risø. 14 out of 41 tested parameters are statistically different with p-level <
30 0.1 (underlined).

31 Supplementary Figure 1. Temporal variation of concentration of selected gases SO_2 , NO, and NO_x
32 (30 min resolution) at HCAB during the campaign period. The x-axis shows a temporal scale from
33 00 - 24h for each day, with the white and grey shades indicating the intervals for day and night
34 samples, respectively.

35 Supplementary Figure 2. Temporal variation (30 min resolution) of the local meteorological
36 parameters at the urban background site HCOE, including relative humidity (RH), global radiation
37 (GR) and temperature (T). The x-axis shows a temporal scale from 00 - 24h for each day, with the
38 white and grey shades indicating the intervals for day and night samples, respectively.

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1 **Figures**

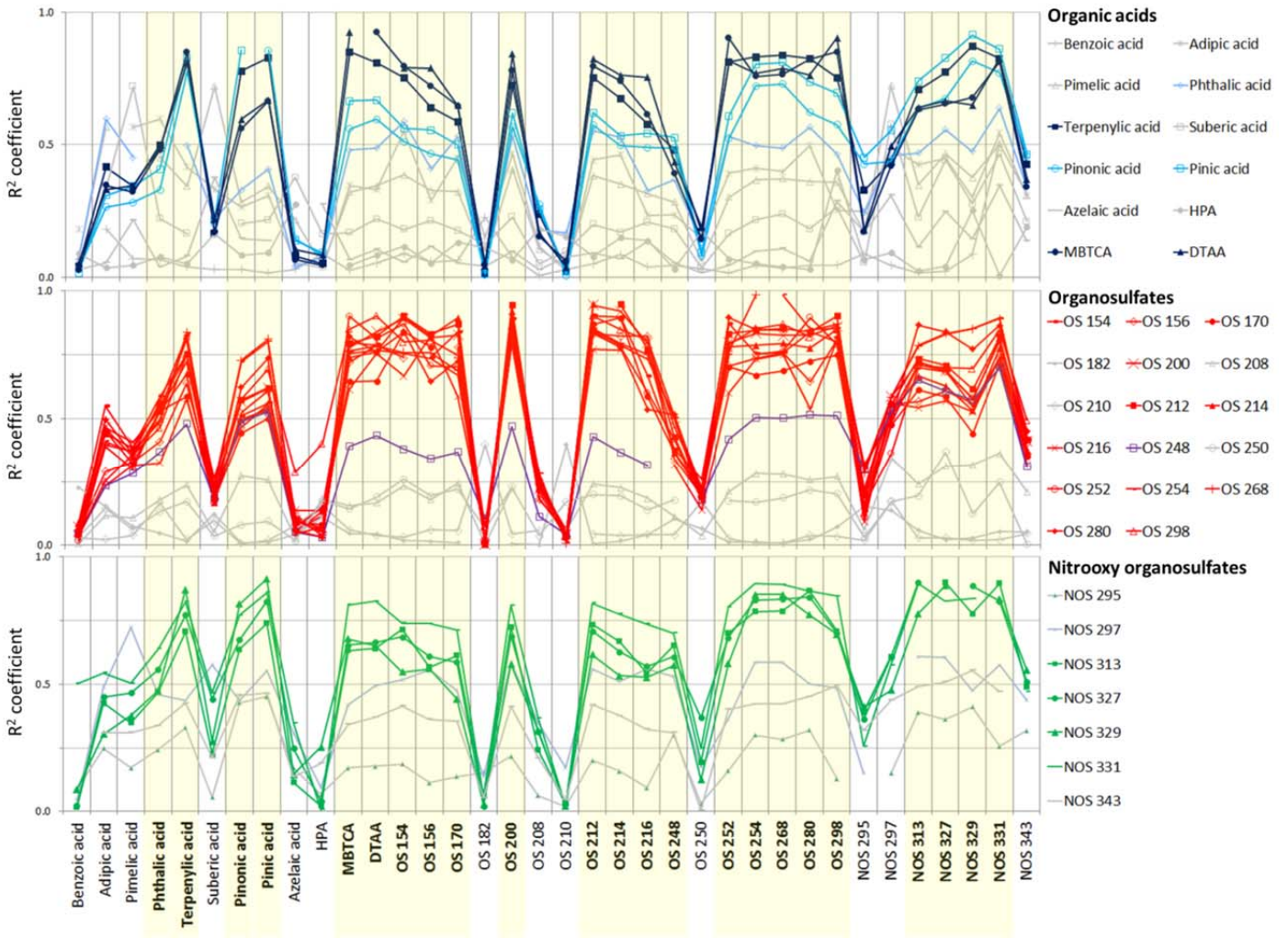
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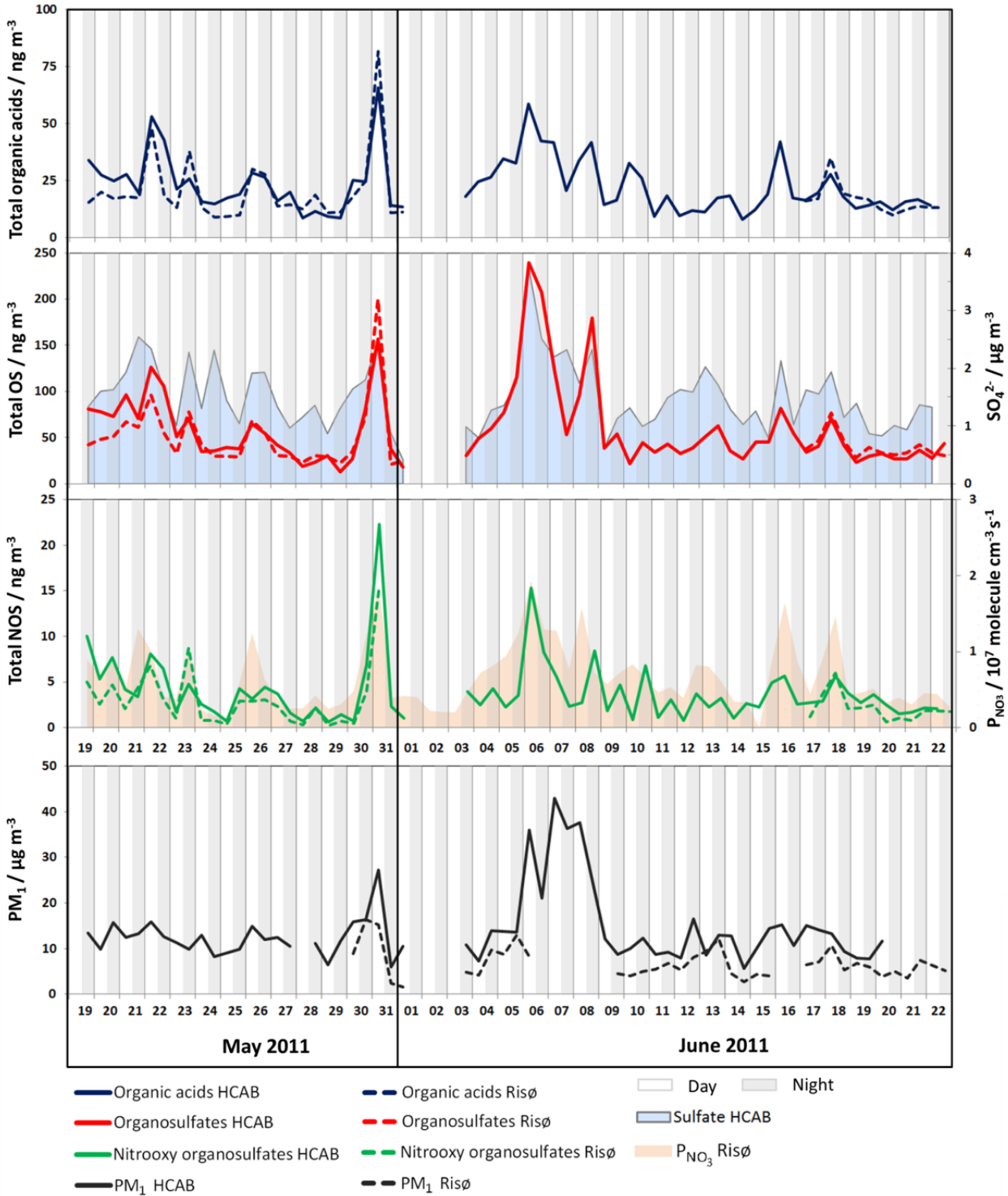
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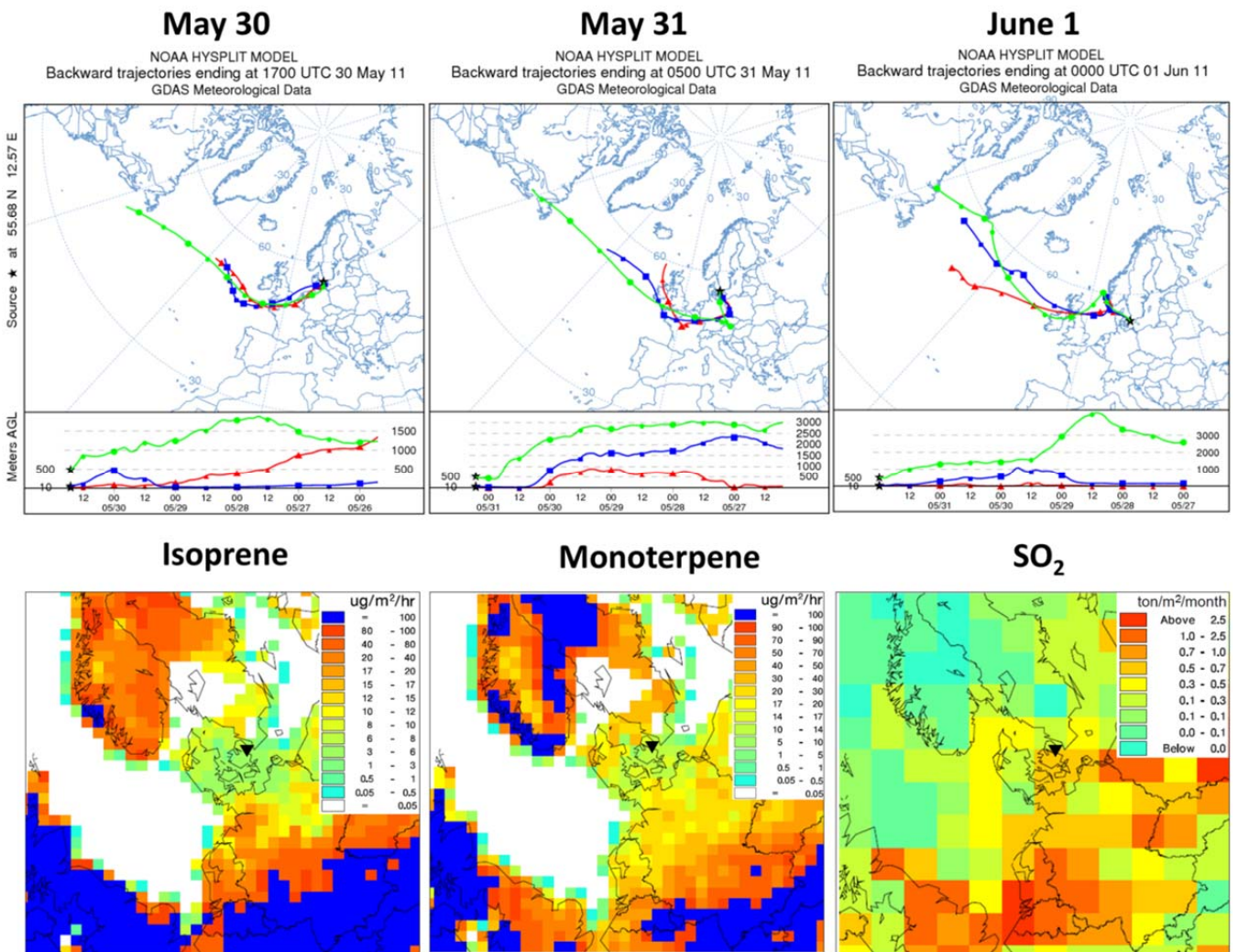


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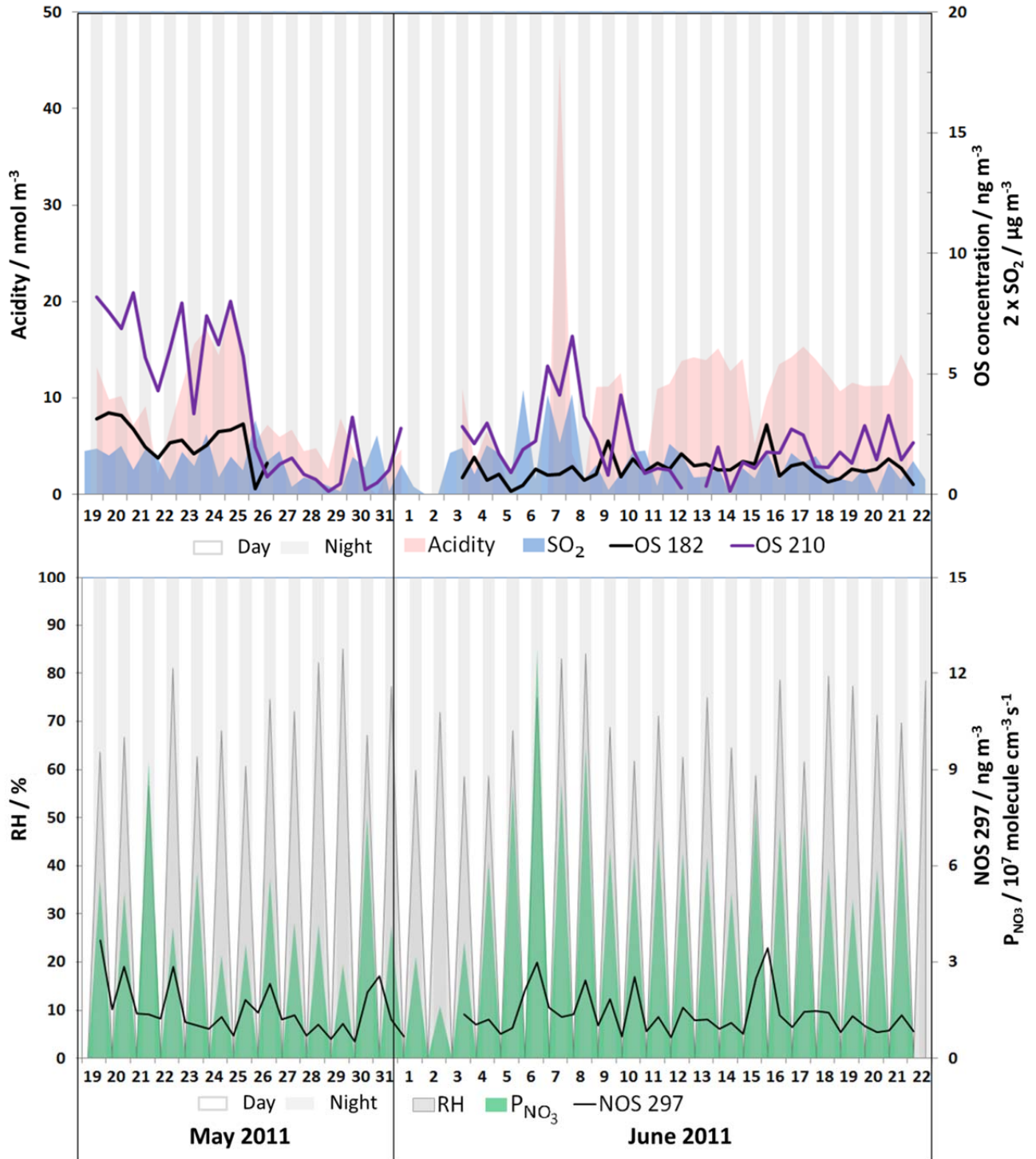


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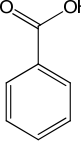
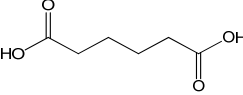
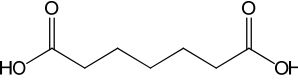
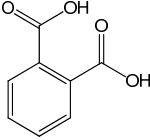
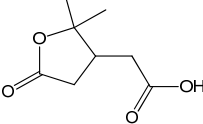
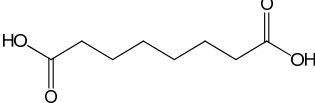
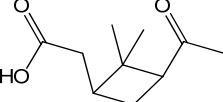
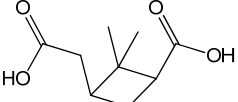
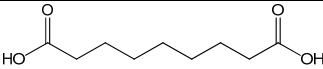
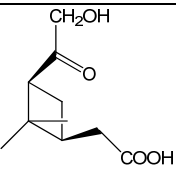
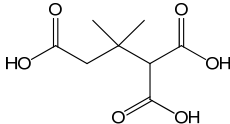
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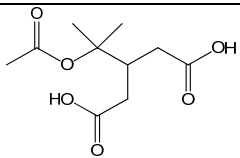
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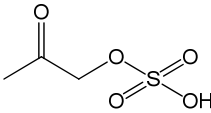
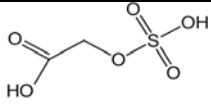
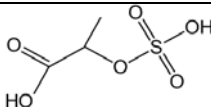
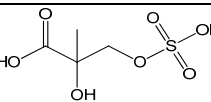
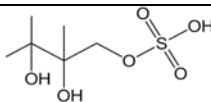
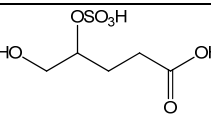
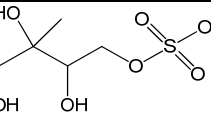
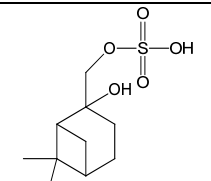
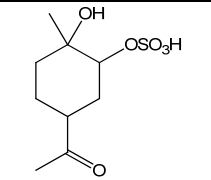
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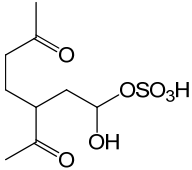
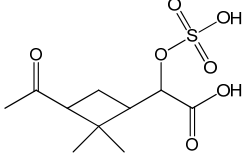
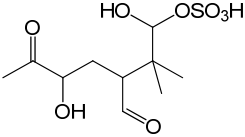
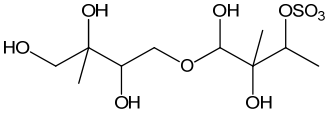
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Organic acid	[M-H] ⁻ ion (m/z)	Retention time (min)	Proposed molecular formula	Proposed structure	Suggested precursors
Benzoic acid	121.022	27.5	C ₇ H ₆ O ₂		Aromatic hydrocarbons
Adipic acid	145.058	14.9	C ₆ H ₁₀ O ₄		Cyclic olefins
Pimelic acid	159.061	20.3	C ₇ H ₁₂ O ₄		Cyclic olefins
Phthalic acid	165.123	19.0	C ₈ H ₆ O ₄		Aromatic hydrocarbons
Terpenylic acid	171.058	18.3	C ₈ H ₁₂ O ₄		α-pinene
Suberic acid	173.074	25.4	C ₈ H ₁₄ O ₄		Fatty acids
Pinonic acid	183.100	27.5	C ₁₀ H ₁₆ O ₃		α/β-pinene
Pinic acid	185.077	23.0; 24.0	C ₉ H ₁₄ O ₄		α/β-pinene
Azelaic acid	187.091	28.1	C ₉ H ₁₆ O ₄		Fatty acids
Hydroxy-pinonic acid	199.092	29.0	C ₁₀ H ₁₆ O ₄		α/β-pinene
3-Methyl-1,2,3-butanetricarboxylic acid MBTCA	203.050	16.2	C ₈ H ₁₂ O ₆		α/β-pinene

Di-terpenylic acid acetate (DTAA)	231.220	24.3	$C_{10}H_{16}O_6$		α -pinene
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1 Table 2. Detected organosulfates (OS) during the campaign.

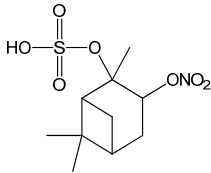
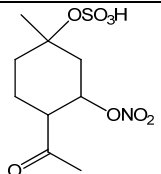
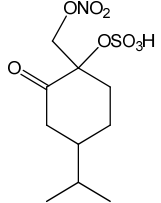
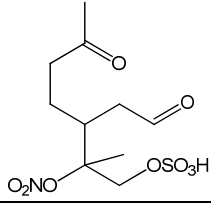
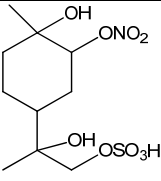
Organosulfate	[M-H] ⁻ ion (m/z)	Retention time (min)	Proposed molecular formula	Proposed structure	Suggested precursors
OS 154 ^{1,2}	152.981	3.6; 6.7	C ₃ H ₆ O ₅ S		hydroxyacetone (isoprene)
OS 156 ³	154.957	2.8	C ₂ H ₄ O ₆ S		glyoxal
OS 170 ⁴	169.047	3.4	C ₃ H ₆ O ₆ S		methylglyoxal
OS 182	181.063	34.8	C ₅ H ₁₀ O ₅ S	unknown	unknown
OS 200_1 ¹	198.998	3.3	C ₄ H ₈ O ₇ S		2-methylglyceric acid (isoprene)*
OS 200_2 ^{5,6}	199.001	3.4	C ₅ H ₁₂ O ₆ S		2-methyl-3-buten-2-ol (MBO)
OS 208	207.101	28.4; 29.5	C ₆ H ₈ O ₆ S	unknown	unknown
OS 210	209.086	28.0; 30.5	C ₇ H ₁₄ O ₅ S	unknown	unknown
OS 212 ²	210.983	4.8	C ₅ H ₈ O ₇ S	unknown	isoprene
				unknown	isoprene ^{2,5}
OS 214 ^{2,7,8}	213.001	3.5	C ₅ H ₁₀ O ₇ S		fatty acid ⁴
OS 216 ^{1,2}	215.018	2.9	C ₅ H ₁₂ O ₇ S		2-methyltetrols (isoprene)**
OS 248 ²	247.061	32.2	C ₁₀ H ₁₆ O ₅ S	unknown	α-pinene
OS 250 ²	249.074	31.6	C ₁₀ H ₁₈ O ₅ S		β-pinene***
OS 252 ²	251.017	15.3	C ₉ H ₁₆ O ₆ S		limonene
OS 254 ²	253.158	21.3	C ₈ H ₁₄ O ₇ S	unknown	α-terpinene

OS 268 ²	267.225	19.7	C ₉ H ₁₆ O ₇ S		limonene
OS 280 ²	279.046	22.5; 25.2	C ₁₀ H ₁₆ O ₇ S		α/β-pinene
OS 298 ²	297.079	25.5	C ₁₀ H ₁₈ O ₈ S		α-pinene
OS 334 ²	333.071	38.0	C ₁₀ H ₂₂ O ₁₀ S		Isoprene

1 Included references were ¹ Surratt et al. (2007a); ² Surratt et al. (2008b); ³ Galloway et al. (2009); ⁴ Olson et al. (2011); ;
2 ⁵ Zhang et al. (2012c) ⁶ Zhang et al (2014) ⁷ Gómez-González et al., (2008); ⁸ Kristensen and Glasius, (2011). * OS 200
3 has been recently shown to specifically form from methacrylic acid epoxide (MAE) uptake onto acidified sulfate
4 aerosol (Lin et al., 2013). ** OS 216 has been shown to specifically derive from the reactive uptake of isoprene
5 epoxydiols (IEPOX) in the presence of acidic sulfate aerosol (Surratt et al., 2010; Lin et al., 2012). *** OS 250 can be
6 found in synthesis of both α and β-pinene derived organosulfates; however with different retention times. The OS 250
7 detected in this study was assigned to β-pinene precursor based on the corresponding retention time.

8

1 Table 3. Detected nitrooxy organosulfates (NOS) during the campaign.

Nitrooxy organosulfate	[M-H] ⁻ ion (m/z)	Retention time (min)	Proposed molecular formula	Proposed structure	Suggested precursors
NOS 295 ²	294.063	42.1; 44.7	C ₁₀ H ₁₇ NO ₇ S	 More isomers	α/β-pinene
NOS 297 ²	296.063	32.8	C ₉ H ₁₅ NO ₈ S		limonene
NOS 311 ²	310.088	38.0	C ₁₀ H ₁₇ NO ₈ S		limonene, α-pinene
NOS 313 ²	312.209	28.0	C ₉ H ₁₅ NO ₉ S	unknown	limonene
NOS 327 ²	326.279	26.8	C ₁₀ H ₁₇ NO ₉ S		limonene, β-pinene, terpinolene
NOS 329 ²	327.994	33.5	C ₁₀ H ₁₉ NO ₉ S		limonene
NOS 331 ²	330.002	15.6	C ₉ H ₁₇ NO ₁₀ S	unknown	unknown
NOS 340 ²	339.070	25.7	C ₁₀ H ₁₆ N ₂ O ₉ S	unknown	α-pinene
NOS 343 ²	342.043	38.8	C ₁₀ H ₁₇ NO ₁₀ S	unknown	α-pinene; α-terpinene

2 References include ¹ Surratt et al., (2007a); ² Surratt et al., (2008b); ³ Surratt et al., (2010); ⁴ Gómez-González et al.,
 3 (2008); ⁵ Kristensen and Glasius, (2011).

4

1 Table 4. Concentration range (ng m⁻³) of the detected species at HCAB and Risø, reported as mean,
 2 standard deviation (stdev.), median, max and the number of detected samples (N) at each site.

Compound	HCAB					Risø				
	Mean	Stdev.	Median	Max	N	Mean	Stdev.	Median	Max	N
Benzoic acid	0.4	0.6	0.1	2.0	14	1.4	1.9	0.4	7.2	26
Adipic acid	3.3	1.4	2.8	8.4	64	2.9	1.1	2.8	5.9	38
Pimelic acid	0.6	0.3	0.5	2.1	64	0.4	0.2	0.4	1.2	38
Phthalic acid	5.6	2.3	5.1	13.0	64	4.3	1.8	4.4	10.4	38
Terpenylic acid	1.6	1.6	1.1	8.9	64	1.4	1.7	0.9	8.5	38
Suberic acid	1.1	0.6	1.0	3.7	64	0.9	0.5	0.8	2.9	38
Pinonic acid	1.3	1.5	0.9	10.8	64	2.2	2.9	1.3	16.4	38
<i>cis</i> -Pinic acid	0.5	0.5	0.4	3.9	64	0.6	0.9	0.4	5.6	37
Azelaic acid	4.0	3.3	3.0	19.6	58	1.6	0.8	1.6	4.3	35
Hydroxy-pinonic acid	0.1	0.1	0.1	0.2	12	0.1	0.0	0.1	0.1	4
MBTCA	4.7	5.7	2.3	23.8	64	3.9	5.1	2.0	24.4	38
DTAA	0.4	0.5	0.2	2.0	64	0.2	0.3	0.1	1.4	34
Total organic acids	22.8	14.9	18.7	76.7	64	19.3	13.3	15.7	69.8	38
OS 154	6.6	6.0	4.5	25.7	63	5.3	4.6	4.3	24.6	38
OS 156	5.3	3.8	4.3	23.1	64	4.3	3.1	3.5	19.8	38
OS 170	3.5	2.3	2.7	10.3	63	3.1	2.0	2.5	11.5	37
OS 182	1.4	0.8	1.2	3.4	55	0.6	0.4	0.6	1.6	37
OS 200_1	7.1	6.7	4.4	33.1	64	4.5	3.9	3.0	22.2	37
OS 200_2	6.4	6.3	4.1	18.4	5	3.1	2.1	2.6	6.7	5
OS 208	5.0	3.3	5.7	14.2	63	4.1	0.8	4.1	5.6	38
OS 210	2.8	2.4	2.0	8.4	63	1.8	1.0	1.9	4.3	38
OS 212	7.4	6.0	5.0	32.2	64	6.2	4.7	4.6	27.8	38
OS 214	5.1	4.5	3.6	22.3	64	4.4	3.1	3.8	17.9	38
OS 216	5.0	6.8	2.8	36.9	60	3.2	3.3	2.4	20.8	34
OS 248	0.8	0.4	0.8	2.3	57	0.6	0.3	0.6	1.7	37
OS 250	2.3	1.1	2.1	6.8	64	2.4	2.1	1.9	12.4	38
OS 252	1.2	1.2	0.6	5.2	53	0.9	0.9	0.6	4.1	33
OS 254	1.3	1.5	0.8	7.9	63	1.1	1.2	0.8	7.4	38
OS 268	2.5	2.3	1.6	13.1	64	2.2	2.2	1.6	13.9	38
OS 280	2.6	2.5	1.8	11.5	60	2.2	2.0	1.6	10.3	38
OS 298	1.3	1.1	0.8	4.1	28	0.8	1.0	0.5	3.9	16
OS 334	0.6	0.3	0.6	1.0	8	0.8	-	0.8	0.8	1
Total organosulfates	59.8	44.5	43.8	130.6	64	46.9	31.1	36.3	144.0	38
NOS 295	0.5	0.6	0.3	4.2	46	0.4	0.3	0.4	1.2	15
NOS 297	1.4	0.7	1.3	3.7	64	0.9	0.6	0.7	3.4	38
NOS 311	0.9	0.8	0.4	2.4	7	0.5	0.5	0.4	1.8	8
NOS 313	0.7	0.7	0.5	3.4	56	0.6	0.6	0.5	3.0	30
NOS 327	0.9	0.7	0.7	3.6	48	0.6	0.6	0.5	2.8	29
NOS 329	0.8	0.9	0.4	3.6	19	0.5	0.6	0.3	2.6	17
NOS 331	0.3	0.2	0.2	1.2	41	0.3	0.3	0.2	1.5	26
NOS 340	0.2	0.1	0.2	0.2	3	-	-	-	-	-
NOS 343	0.8	0.4	0.8	1.4	24	0.4	0.4	0.4	1.0	4
Total nitrooxy organosulfates	3.9	1.3	2.8	3.7	64	2.8	1.2	2.1	1.8	36